## Figure 1 125P5C8 SSH sequence

Series Marie Marie

Harle Same and Start

Start II II II'' Ama Haif

Figure 2 Double stranded sequence and ORF for 125P5C8-Pro-pCR2.1.

	М	Т	s	L	M	R	E	I	L	L	E	S	L	L	G	С	V	S
1	ATG W	ACC S	TCG L	CTG Y	TGG H	AGA D	GAA L	ATC G	CTC P	TTG M	GAG I	TCG Y	CTG Y	CTG F	GGA P	TGT L	GTT Q	TCT
55		-		_			_	GGA					TAC			TTG	CAA	
100	L	E	L	T	G	L CTT	E GAA	G GGT	F TTT	S AGT	I ATA	A GCA	$_{ m TTT}$	L CTT	S TCT	P CCA	I ATA	F TTC
109	L	GAA T	CTC	AC1 T	GGG P	F	W	K	Ŀ	V	N	K	K	M	M	L	Т	L
163	CTA	ACA		ACT								AAG		TGG	ATG P	CTA N	_	CTG K
217	L CTG	R AGG	I ATA	I	T ACT	I ATT	G GGC	S AGC	I ATA	A GCC	S TCC	F TTC	Q CAG	A GCT	CCA		A GCC	AAA
22,	L	R	L	М	V	L	A	$\mathbf{L}$	G	V	S	S	S	L	I	V	Q	A
271	CTT V	CGA T	CTG W	ATG W	GTT S	CTT	GCG S	CTT H	GGG L	GTG Q	TCT R	TCC Y	TCA L	CTG R	ATA I	GT.G	CAA G	GCT F
325	GTG	ACT		TGG		GGA	-	CAT		CAA		TAC		AGA	ATT	TGG	GGA	TTC
	I	L	G	Q	I	V	L	V	V	L	R	I ATA	W TCC	Υ mvm	T ACT	S TCA	L CTA	N AAC
379	ATT P	TTA	GGA W	CAG S	Y	GTT O	CTT M	S	GTT N	K	V	I	L	T	L	Ş	A	I
433	CCA	ATC	TGG	AGT	TAT							ATA				AGT	GCC	ATA
487	A GCC	T ACA	L CTT	D GAT	R CGT	I ATT	GGC	T ACA	D GAT	G GGT	D GAC	C TGC	S AGT	K AAA	P CCT	E GAA	E GAA	K AAG
407	K	T	G	E	V	A	Т	G	M	A	S	R	P	N	W	$\mathbf{L}$	L	Α
541		ACT	GGT	GAG		GCC		GGG V	ATG F	GCC L	TCT T	AGA H	CCC W	AAC V	TGG F	CTG G	CTG E	GCA V
595	G GGG	A GCT	A GCT	F TTT	G GGT	S AGC	L CTT	or GTG		CTC			TGG		TTT	GGA	_	GTC
	S	${f L}$	V	S	R	W	A	V	S	G	Н	P	Н	P	G	P	D	P
649	TCT N	CTT P	GTT F	TCC G	AGA G	TGG A	GCA V	GTG L	AGT L	GGG C	CAT	CCA A	CAT	CCA G	GGG L	M	GAT L	CCT P
703	AAC	CCA	TTT	GGA	GGT	GCA		CTG		TGC	TTG	GCA		GGA	TTG		CTT	CCA
252	S	С	L	W TGG	F TTT	R CGT	G COT	T ACT	G GGT	L TTG	I ATC	W TGG	W TGG	ÇTTT V	T ACA	G GGA	T	A GCT
757	TCT S	TGT A	TTG A	G	T T T	L CG1	Y	L	H	T	W	A	A	A	V	S	G	C
811	TCA			GGG	CTC	CTT	TAC					GCA		GCT	GTG G	TCT H	GGC L	TGT I
865	V GTC	F TTC	A GCC	I ATC	F TTT	T ACT	A GCA	S TCC	M ATG	W TGG	P CCC	Q CAA	T ACA	L CTT	GGA	CAC	CTT	ATT
303	N	s	G	Т	N	P	G	K	Т	M	$\mathbf{T}$	I	A	M	I	_F_	Y	L
919	AAC L	TCA E	GGG I	ACA F	AAC F	CCT	GGG A	AAA W	ACC C	ATG T	ACC A	ATT F	GCC K	ATG F	ATA V	TTT P	TAT G	CTT G
973	CTA		ATA	_	TTC	TGT	GCC			ACA		TTT	AAG	TTT	GTC		GGA	GGT
	V	Y	A	R	E	R	S	D	V	L	L TTG	G	T	M	M ATG	L	I ATT	I ATC
1027	GTC G	TAC L	GCT N	AGA M	GAA L	AGA F	TCA G	GAT P	GTG K	CTT K	N	L	D	L	L	L	Q	T
1081	GGG		AAT		CTA		GGT			AAA				TTG		CTT	CAA	
1135	K AAA	N AAC	S agt	S TCT	K AAA	V GTG	$_{ m CTT}$	F TTC	R AGA	K AAG	S AGT	E GAA	K AAA	Y TAC	M ATG	K AAA	L CTT	${ m F} \ { m TTT}$
1133	L	W	$_{ m L}$	L	V	G	V	G	L	L	G	$_{ m L}$	G	$\mathbf{L}$	R	H	K	A
1189	CTG Y	TGG E	CTG R	CTT K	GTT L	GGT G	GTG K	GGA V	TTG A	TTG P	GGA T	TTA K	GGA E	CTA V	CGG S	CAT A	AAA A	GCC
1243	TAT														TCT			
	W	P	F	R	F	G	Y	D	N	E	G	W	S	S	L	E	R	S
1297	TGG A	H CCL	TTC	AGG L	N L.T.T.	GGA E	TAT	GAC G	AAT	GAA D	F	I	T	I	CTA L	E	S	D
1351	GCT		CTG	CTC											TTG			
1405	A CCT	S TCT	K NAG	P	Y TAT	M ATG	G	N AAC	N AAT	D GAC	L ATT	T ACC	M ATG	W TGG	L CTA	G GGG	E GAA	K AAG
1405	$\mathbf{L}$	G	F	Y	$\mathbf{T}$	D	F	G	P	S	T	R	Y	H	Т	W	G	I
1459		GGT								AGC K	ACA S	AGG E	TAT H	CAC H	ACT L	TGG L	GGG P	ATT S
1513	M ATG	A GCT	L TTG	S TCA	R AGA	Y TAC	P CCA	I . ATT	V GTG									
	P	E	G	E	I	Α	P	A	I	Т	L	T	$\Lambda$	N	I	S	G	K
1567	CCA L	GAG V	GGC D	GAG F	ATC V	GCA V	. CCA T	GCC H	ATC F	ACA G	. TTG N	ACC H	GTT E	AAC D	ATT D	TCG L	GGC D	AAG R

## Figure 2 Cont'd

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1621	CTG	GTG	GAT	TTT	GTC	GTG	ACA	CAC	TTT	GGG	AAC	CAC	GAA	${\tt GAT}$	GAC	CTC	GAC	AGG
	K	L	Q	A	I	A	V	S	K	L	$_{ m L}$	K	S	S	S	N	Q	V
1675	AAA	CTG	CAG	GCT	ATT	GCT	GTT	TCA	AAA	CTA	CTG	AAA	AGT	AGC	TCT	TAA	CAA	GTG
	I	F	L	G	Y	I	$\mathbf{T}$	S	Α	P	G	S	R	D	Y	L	Q	$\mathbf{L}$
1729	ATA	TTT	CTG	GGA	TAT	ATC	ACT	TCA	GCA	CCT	GGC	TCC	AGA	GAT	TAT	CTA	CAG	CTC
	${f T}$	E	H	G	N	V	K	D	Ι	D	S	T	D	Η	D	R	M	C
1783	ACT	GAA	CAT	GGC	AAT	GTG	AAG	GAT	$\operatorname{ATC}$	GAC	AGC	ACT	GAT	CAT	GAC	AGA	TGG	TGT
	E	Y	I	M	Y	R	G	L	I	R	L	G	Y	Α	R	I	S	H
1837	GAA	TAC	ATT	ATG	TAT	CGA	GGG	CTG	${\tt ATC}$	AGG	TTG	GGT	TAT	GCA	AGA	ATC	TCC	CAT
	A	E	L	S	D	S	E	I	Q	M	A	K	F	R	I	P	D	D
1891	GCT	GAA	CTG	AGT	GAT	TCA	GAA	$\operatorname{ATT}$	CAG	ATG	GCA	AAA	TTT	AGG	ATC	CCT	GAT	GAC
	P	$\mathbf{T}$	N	Y	R	D	N	Q	K	V	V	I	D	Η	R	E	V	S
1945	CCC	ACT	AAT	TAT	AGA	GAC	AAC	CAG	AAA	GTG	GTC	ATA	GAC	CAC	AGA	GAA	GTT	TCT
	E	K	I	H	F	N	P	R	F	G	S	Y	K	E	G	Η	N	Y
1999	GAG	AAA	ATT	CAT	TTT	AAT	CCC	AGA	TTT	GGA	TCC	TAC	AAA	GAA	GGA	CAC	AAT	TAT
	E	N	N	H	H	F	H	M	N	T	P	K	Y	F	L	*		
2053	GAA	AAC	AAC	CAT	CAT	TTT	CAT	ATG	AAT	ACT	CCC	AAA	TAC	TTT	TTA	TGA	AAC	

Figure 3 Amino acid sequence of 125P5C8.

1	MTSLWREILL	ESLLGCVSWS	LYHDLGPMIY	YFPLQTLELT	GLEGFSIAFL	50
51	SPIFLTITPF	WKLVNKKWML	TLLRIITIGS	IASFQAPNAK	LRLMVLALGV	100
101	SSSLIVQAVT	WWSGSHLQRY	LRIWGFILGQ	IVLVVLRIWY	TSLNPIWSYQ	150
151	MSNKVILTLS	AIATLDRIGT	DGDCSKPEEK	KTGEVATGMA	SRPNWLLAGA	200
201	AFGSLVFLTH	WVFGEVSLVS	RWAVSGHPHP	GPDPNPFGGA	VLLCLASGLM	250
251	LPSCLWFRGT	GLIWWVTGTA	SAAGLLYLHT	WAAAVSGCVF	AIFTASMWPQ	300
301	TLGHLINSGT	NPGKTMTIAM	IFYLLEIFFC	AWCTAFKFVP	GGVYARERSD	350
351	VILGTMMLII	GLNMLFGPKK	NLDLLLQTKN	SSKVLFRKSE	KYMKLFLWLL	400
401	VGVGLLGLGL	RHKAYERKLG	KVAPTKEVSA	AIWPFRFGYD	NEGWSSLERS	450
451	AHLLNETGAD	FITILESDAS	KPYMGNNDLT	MWLGEKLGFY	TDFGPSTRYH	500
501	TWGIMALSRY	PIVKSEHHLL	PSPEGEIAPA	ITLTVNISGK	LVDFVVTHFG	550
551	NHEDDLDRKL	QAIAVSKLLK	SSSNOVIFLG	YITSAPGSRD	YLQLTEHGNV	600
601	KDIDSTDHDR	WCEYIMYRGL	IRLGYARISH	AELSDSEIQM	AKFRIPDDPT	650
	NYRDNOKVVI	DHREVSEKIH	FNPRFGSYKE	GHNYENNHHF	HMNTPKYFL	699
651	MIKDMÖKAAT	DITTE A DELETT	I IVI KI OD I KE	0111, 4 221111111		

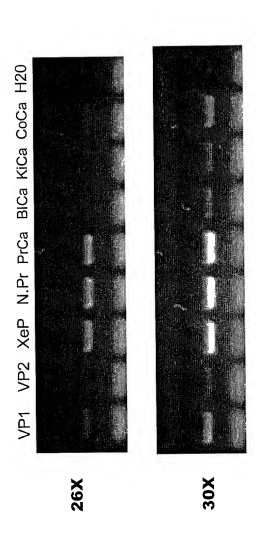
Figure 4A Alignment with AK025164 protein product Score = 1397 bits (3615), Expect = 0.0 Identities = 682/699 (97%), Positives = 683/699 (97%)

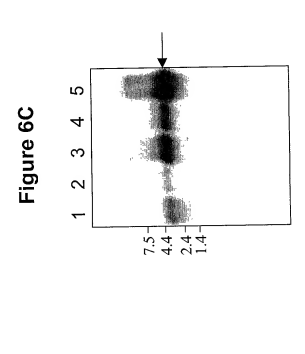
Query:	1	MTSLWREILLESLLGCVSWSLYHDLGPMIYYFPLQTLELTGLEGFSIAFLSPIFLTITPF MTSLWREILLESLLGCVSWSLYHDLGPMIYYFPLQTLELTGLEGFSIAFLSPIFLTITPF	60
Sbjct:	1	MTSLWREILLESLLGCVSWSLYHDLGPMIYYFPLQTLELTGLEGFSIAFLSPIFLTITPF	60
Query:	61	WKLVNKKWMLTLLRIITIGSIASFQAPNAKLRLMVLALGVSSSLIVQAVTWWSGSHLQRY WKLVNKKWMLTLLRIITIGSIASFQAPNAKLRLMVLALGVSSSLIVQAVTWWSGSHLQRY	120
Sbjct:		WKLVNKKWMLTLLRIITIGSIASFQAPNAKLRLMVLALGVSSSLIVQAVTWWSGSHLQRY	
		LRIWGFILGQIVLVVLRIWYTSLNPIWSYQMSNKVILTLSAIATLDRIGTDGDCSKPEEK LRIWGFILGQIVLVVLRIWYTSLNPIWSYQMSNKVILTLSAIATLDRIGTDGDCSKPEEK	
_		LRIWGFILGQIVLVVLRIWYTSLNPIWSYQMSNKVILTLSAIATLDRIGTDGDCSKPEEK	
		KTGEVATGMASRPNWLLAGAAFGSLVFLTHWVFGEVSLVSRWAVSGHPHPGPDPNPFGGA KTGEVATGMASRPNWLLAGAAFGSLVFLTHWVFGEVSLVSRWAVSGHPHPGPDPNPFGGA	
_		KTGEVATGMASRPNWLLAGAAFGSLVFLTHWVFGEVSLVSRWAVSGHPHPGPDPNPFGGA	300
_		VLLCLASGLMLPSCLWFRGTGLIWWVTGTASAAGLLYLHTWAAAVSGCVFAIFTASMWPQ VLLCLASGLMLPSCLWFRGTGLIWWVTGTASAAGLLYLHTWAAAVSGCVFAIFTASMWPQ	
-		VLLCLASGLMLPSCLWFRGTGLIWWVTGTASAAGLLYLHTWAAAVSGCVFAIFTASMWPQ	
_		TLGHLINSGTNPGKTMTIAMIFYLLEIFFCAWCTAFKFVPGGVYARERSDVLLGTMMLII TLGHLINSGTNPGKTMTIAMIFYLLEIFFCAWCTAFKFVPGGVYARERSDVLLGTMMLII TLGHLINSGTNPGKTMTIAMIFYLLEIFFCAWCTAFKFVPGGVYARERSDVLLGTMMLII	
_		GLNMLFGPKKNLDLLLQTKNSSKVLFRKSEKYMKXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX	
_		GLNMLFGPKKNLDLLLQTKNSSKVLFRKSEKYMK RHKAYERKLG GLNMLFGPKKNLDLLLQTKNSSKVLFRKSEKYMKLFLWLLVGVGLLGLGLRHKAYERKLG	
_		KVAPTKEVSAAIWPFRFGYDNEGWSSLERSAHLLNETGADFITILESDASKPYMGNNDLT	
		KVAPTKEVSAAIWPFRFGYDNEGWSSLERSAHLLNETGADFITILESDASKPYMGNNDLT KVAPTKEVSAAIWPFRFGYDNEGWSSLERSAHLLNETGADFITILESDASKPYMGNNDLT	
_		MWLGEKLGFYTDFGPSTRYHTWGIMALSRYPIVKSEHHLLPSPEGEIAPAITLTVNISGK	540
Sbjct:	481	MWLGEKLGFYTDFGPSTRYHTWGIMALSRYPIVKSEHHLLPSPEGEIAPAITLTVNISGK MWLGEKLGFYTDFGPSTRYHTWGIMALSRYPIVKSEHHLLPSPEGEIAPAITLTVNISGK	540
Query:	541	LVDFVVTHFGNHEDDLDRKLQAIAVSKLLKSSSNQVIFLGYITSAPGSRDYLQLTEHGNV	600
Sbjct:	541	LVDFVVTHFGNHEDDLDRKLQAIAVSKLLKSSSNQVIFLGYITSAPGSRDYLQLTEHGNV LVDFVVTHFGNHEDDLDRKLQAIAVSKLLKSSSNQVIFLGYITSAPGSRDYLQLTEHGNV	600
Query:	601	KDIDSTDHDRWCEYIMYRGLIRLGYARISHAELSDSEIQMAKFRIPDDPTNYRDNQKVVI KDIDSTDHDRWCEYIMYRGLIRLGYARISHAELSDSEIQMAKFRIPDDPTNYRDNQKVVI	660
Sbjct:	601	KDIDSTDHDRWCEYIMYRGLIRLGYARISHAELSDSEIQMAKFRIPDDPTNYRDNQKVVI	660
Query:	661	DHREVSEKIHFNPRFGSYKEGHNYENNHHFHMNTPKYFL 699 DHREVSEKIHFNPRFGSYKEGHNYENNH+FHMNTPKYFL	
Sbjct:	661	DHREVSEKIHFNPRFGSYKEGHNYENNHNFHMNTPKYFL 699	

## Figure 4B Alignment with yeast YCR017 cp Protein

Score = 261 bits (668), Expect = 1e-68 Identities = 204/705 (28%), Positives = 330/705 (45%), Gaps = 50/705 (7%) Query: 15 GCVSWSLYHDLGPMIYYFPLQTLELTGLEGFSIAFLSPIFLTITPFWKLVNKKWMLTLLR 74 G + WS L I++FPL + ++G E + +L PIFL + PFSbjct: 279 GFLFWSNVTSLLCSIWHFPLWYMGISGYEAAILGYLGPIFLYL-PFVSEAFTQYGVLLGG 337 Query: 75 IITIGSIASFQAPNAKLRLMVLALGVSSSLIVQAVTWWSGSHLQ-RYLRIWGFILGQIVL 133 Q P + L + + + + + VQ + + + + + W + LG +TT TG+ Sbjct: 338 IIAIGAYI-VQMPELRLISVAVGTSITVATFVQNLRYITNAETSFSFALTW--LLGLVAS 394 Query: 134 VVLRIWYTSLNPIWSYQMS----NKVILTLSAIATLDRIGTDGDCSKPEEKKTGEVATG 188 V+L++ + + NP W NK L L+ + + + E K+ + + Sbjct: 395 VILKMGFYTNNPTWVILDERNGGYNKTALVLTVLFGM--LSPYVNSINFEGKRNAQAKS- 451 Query: 189 MASRPNWLLAGAAFGSLVFLTHWVFGEVSLVSRWAVSGHPHP-GPDPNPFGGAVLLCLAS 247 FGSL+F H + + S WA G+ GP P P+G AS L Sbjct: 452 -ASLIGKLFLAVGFGSLLFGIHQLLTDSSTTIYWAWEGYNESHGPLPWPWGA--LTCTVM 508 Query: 248 GLMLPSCLWFRGTGLIWWVTGTASAAGLLY--LHTWAAAV-SGCVFAIFTASMWPQ---T 301 S + F G L + + S A L + W + G + + AISbjct: 509 LFASLSSVKFMGKPLVPCLLLLISTAVLSARSITQWPKYIFGGLLYAIAMLWLVPSYFSA 568 Query: 302 LGHLINSGTNPGKTMTIAMIFYLLEIFFCAWCTAFKFVPGGVYARERSDVLLGTMMLIIG 361 LG + N ++ Y++ + W A+ FVP G RE+ + +L Sbjct: 569 LGQVQNIWV-----YVLSFSVYIIFVLAHVWVVAYAFVPMGWVLREKIETVLAFSSTFII 623 Query: 362 LNMLFGPKKNLDLLLQTKNSSKVLFRKSEKYMKXXXXXXXXXXXXXXXXRHKAYERKLGK 421 N+ L+ K +F + L Sbjct: 624 IGALTCKNLNIQLVTMGKKFFIYVF------FFAVALLSLTARFVYDIRPTGI 670 Query: 422 VAP----TKEVSAAIWPFRFGYDNEGWSSLERSAHLLNETGADFITILESDASKPYMGNN 477 ++ ++A IW FG DN+ W+S +R +L+ + D + +LE+D + MGN Р Sbjct: 671 PQPYHPDSQLITAGIWTIHFGLDNDMWASEDRMINLIKDMELDVVGLLETDTQRITMGNR 730 Query: 478 DLTMWLGEKLGFYTDFGPSTRYHTWGIMALSRYPIVKSEHHLLPSPEGEIAPAITLTV-N 536 DLT L L Y DFGP HTWG + LS++PIV S HHLLPSP GE+APAI T+ Sbjct: 731 DLTSKLAHDLNMYADFGPGPNKHTWGCVLLSKFPIVNSTHHLLPSPVGELAPAIHATLQT 790 Query: 537 ISGKLVDFVVTHFGNHEDDLDRKLQAIAVSKLLKSSSNQVIFLGYITSAPGSRDY-LQLT 595 + LVD V H G ED+ DR+LQ+ ++KL+ +++ I L Y+ PG +Y Sbjct: 791 YNDTLVDVFVFHSGQEEDEEDRRLQSNYMAKLMGNTTRPAILLSYLVVDPGEGNYNTYVS 850 Query: 596 EHGNVKDIDSTDHDRWCEYIMYRGLIRLGYARISHAELSDSEIQMAKFRIPDDPTNYRDN 655 + DID +D DRWCEYI+YRGL R GYAR++ ++D+E+Q+ KF++ + ++ Sbjct: 851 ETSGMHDIDPSDDDRWCEYILYRGLRRTGYARVARGTITDTELQVGKFQVLSEQA-LVEH 909 Query: 656 QKVVIDHREVSEKIHFNPRFGSYKEGHNYENNHHFHM-NTPKYFL 699 + ++ +SE + + +F G E H +H+ + P+Y+L Sbjct: 910 SDSMYEYGHMSEPEYEDMKFPDKFLGEG-ERGHFYHVFDEPRYYL 953

Figure 5





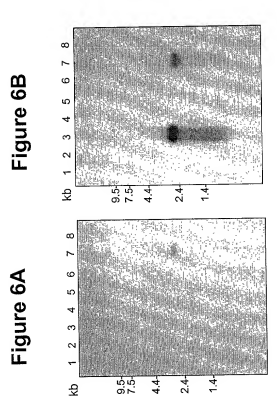


Figure 7

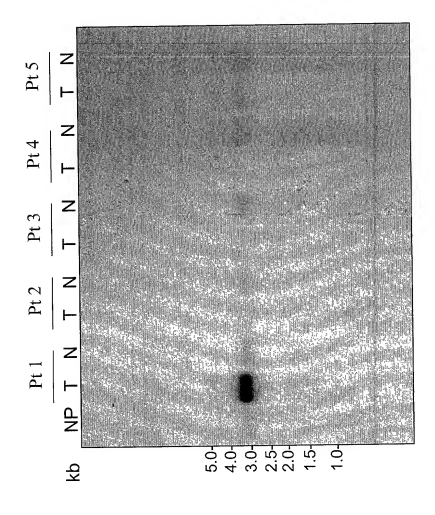


Figure 8

